#### APRIL/MAY 2023

### **CBT53** — BIOINFORMATICS

Time: Three hours

Maximum: 75 marks

LIBRARY

SECTION A —  $(10 \times 2 = 20 \text{ marks})$ 

Answer ALL questions.

- What are the uses of NCBI?
- 2. Summarize about specialized databases.
- 3. Recall the formula of Needleman wunsch algorithm.
- 4. Compare Global and local alignment with examples.
- 5. Name any three gene prediction methods.
- 6. Outline the significance of comparative genomics.
- 7. List out the features of Rasmol.
- 8. Compare SCOP and CATH.
- 9. Define comparative modeling.
- 10. Summarize about alpha helix.

## SECTION B — $(5 \times 5 = 25 \text{ marks})$

### Answer ALL questions.

11. (a) Organize the applications of Bio informatics in various fields.

Or

- (b) List out the protein primary databases and explain.
- 12. (a) Utilize progressive alignment for phylogenetic tree construction.

Or

- (b) Construct the steps in dot plot method to compare two sequences with an example.
- 13. (a) Categorize the steps in performing FASTA.

Or

- (b) List out any three genomic analysis tools and explain any one.
- 14. (a) Identify the features of protein structure database.

Or

- (b) Categorize the applications of protein visualization tools.
- 15. (a) How will you utilize the tools to predict protein functions.

Or

(b) Compare Steps in ab initio modelling and fold recognition method.

# SECTION C $-(3 \times 10 = 30 \text{ marks})$

Answer any THREE questions.

16. Explain in detail about Biological databases.



- . Compare pair wise and multiple sequence alignment.
  - Explain the steps of BLAST algorithm and its applications.
- 19. Elaborate in detail protein classification based on structure.
- 20. Discuss in detail about steps in homology modeling.